⁵^T ANT (GETTTTCCACGTTTTGCNTGACCCTGTTTGCTCAACTRWCKTYTKTKTYKYKTTYTSTKTTRYGCSSYKWYAMRAKMYM SECR IDIN : 1 MRMKTTKAAAAAMCMRRAAAGTTAAYTGGTAAGTTTAGTCTTTTTGTCTTTTATTTCAAGGTCCCGGATCCGGTGGTGG TGCAAATCAAAGAACTGCTCCTCAGTGGATGTTGCCTTTACTTCTAGGCCTGTACGGAAGTGTTACTTCTGCTCTAAAA GCTGCGGAATTCTAATACGACTCACTATAGGGAGTCGACCCACGCGTCCGGTGGGCAGGCCGGGGTGAGGCCTCGCGC TCCGGGAGCTGCACGGGGCTGCGTGGAAAGAGCGCCGAGCGGTCGTTGTCGCCCCCTCCTCGTCGGGAAGAATC START SEDIDNU Z M P SR T Q L F F P T. I R N CEL I Y CCA ACT CAA CTG TTT TTT CCT CTC ATC CGT AAC TGT GAA CTG AGC AGG ATC TAT GGC ACT 43 Y C Н H K H L С C S S S Y I GCA TGT TAC TGC CAC CAC AAA CAT CTC TGT TGT TCC TCA TCG TAC ATT CCT CAG AGT CGA 129 63 Y T P H P A Y Α T F С R P K E N W CTG AGA TAC ACA CCT CAT CCA GCA TAT GCT ACC TTT TGC AGG CCA AAG GAG AAC TGG TGG 189 T Q G R R Y A **S** Т P Q K F Y L T 83 CAG TAC ACC CAA GGA AGG AGA TAT GCT TCC ACA CCA CAG AAA TTT TAC CTC ACA CCT CCA QUV N S I L K A N 103 E Y SFKVP E F CAA TTC AAT AGC ATC CTT AAA GCT AAT GAA TAC AGT TTC AAA GTG CCA GAA TTT GAC GGC 309 K N V S S I L G F D SN QLPAN 123 AAA AAT GTC AGT TCT ATC CTT GGA TTT GAC AGC AAT CAG CTG CCT GCA AAT GCA CCC ATT 369 E DRR TCLQTRGMLL 143 SAA G GAG GAC COG AGA AGT GCA GCA ACC TGC TTG CAG ACC AGA GGG ATG CTT TTG GGG GTT TTT 429 D G H A G C A C S Q A V S E R L GAT THE CAT GCA GGT TGT GCT TGT TCC CAG GCA GTC AGT GAA AGA CTC TTT TAT TAT ATT S L L P H E T L L E I E N A v F. 183 GCT GTC TCT TTG TTA CCC CAT GAG ACT TTG CTA GAG ATT GAA AAT GCA GTG GAG AGC GGC 549 S 203 I RALLP L Q W H K H P N D Y F CGG GCA CTG CTA CCC ATT CTC CAG TGG CAC AAG CAC CCC AAT GAT TAC TTT AGT AAG GAG W F N S T Y Q E L I D. K L Y L R GCA TCC AAA TTG TAC TTT AAC AGC TTG AGG ACT TAC TGG CAA GAG CTT ATA GAC CTC AAC T E S Т D I D v K E A L I N Α F K 243 ACT GGT GAG TCG ACT GAT ATT GAT GTT AAG GAG GCT CTA ATT AAT GCC TTC AAG AGG CTT 263 S L E A Q V G D P N S F N I GAT AAT GAC ATC TCC TTG GAG GCG CAA GTT GGT GAT CCT AAT TCT TTT CTC AAC TAC CTG 789 S T D 283 R Α F G Α A C V Α H GTG CTT CGA GTG GCA TTT TCT GGA GCC ACT GCT TGT GTG GCC CAT GTG GAT GGT GTT GAC 849 T G D S R v Q 303 A M L G CTT CAT GTG GCC AAT ACT GGC GAT AGC AGA GCC ATG CTG GGT GTG CAG GAA GAG GAC GGC 909

Figure 1A

323 A V Т D HNAQNE R F. TCA TGG TCA GCA GTC ACG CTG TCT AAT GAC CAC AAT GCT CAA AAT GAA AGA GAA CTA GAA 969 D 343 РК V V K Q R K L E H S E K S Α CGG CTG AAA TTG GAA CAT CCA AAG AGT GAG GCC AAG AGT GTC GTG AAA CAG GAT CGG CTG 1029 M P F R F G D V K F ĸ 363 A 1089 CTT GGC TTG CTG ATG CCA TTT AGG GCA TTT GGA GAT GTA AAG TTC AAA TGG AGC ATT GAC I E S G ₽ D Q L N D N 383 CTT CAA AAG AGA GTG ATA GAA TCT GGC CCA GAC CAG TTG AAT GAC AAT GAA TAT ACC AAG 1149 N Y H T P P Y L T E E v 403 P 1209 423 R P Q D K F L Λ. L Α т D G L W CAC CGA TTA AGG CCA CAG GAT AAG TTT CTG GTG TTG GCT ACT GAT GGG TTG TGG GAG ACT 1269 V R R Q D v I V G E Y L T G M H 443 ATG CAT AGG CAG GAT GTG GTT AGG ATT GTG GGT GAG TAC CTA ACT GGC ATG CAT CAC CAA 1329 V G G Y K V T G Q M H G 463 L CAG CCA ATA GCT GTT GGT GGC TAC AAG GTG ACT CTG GGA CAG ATG CAT GGC CTT TTA ACA 1389 E R R T K M S s v F E D Q N A A T H I 483 GAA TAGG AGA ACC AAA ATG TCC TCG GTA TTT GAG GAT CAG AAC GCA GCA ACC CAT CTC ATT V G N 503 R : I PRE H Α N E F G т v D Н E R L S CGC CAC GCT GTG GGC AAC AAC GAG TTT GGG ACT GTT GAT CAT GAG CGC CTC TCT AAA ATG 1509 LUSLPEELARM Y R D D I T I 523 CTT AGT CTT CCT GAA GAG CTT GCT CGA ATG TAC AGA GAT GAC ATT ACA ATC ATT GTA GTT N Q B * 1569 Q F N S H V V G Α Y Q 538 CAGITTC AAT TOT CAT GTT GTA GGG GCG TAT CAA AAC CAA GAA TAG End of SEQ IDNO: 3 1614 TGAÇÇGCTCTTTCACTGGCAATTCTCAAATGATATACATTTAAAGGGCAGATTTTTAAAAAGATACTACTATAATAA GGFEGCAGGGTCAGGAGAGTCTGGTCCTGCCTAGCTCAGATTTCATGGCACCTGCACTTGAAGCAAGTCACTTCTTTA TCACAGGTGTCTTGAAACATTAGCTTCTTTTACCAACCTGAGAAAATTAGGATGACCTGGCAAATAAGATCTTGAATAG GCCAAAAGCAAGTATCTTGCTGTGTGTGTCTCTTGGTTAAAGTGAAGAAACAGTACTGTTCACACCTTTCTTCACTGA GATTCCAGTGTACATGAGAACATATATTTATTKSMWKRWTTTYYWRRTACACAGTCTATGCATTWTTCATAWWMAWTTA TTTTWGCCTAAATAARGTKKTTWWCAMATCYAGTTHWTCMATCMATRAACRASMAMCAASCAATCTRTATKTRTTTTTK TKWKTRWTTRWYTGRMAKGMWTSYTWAKTRRAKRAMTAWMCWCMSTYATCCAYCCGMYYKMYTWMYKWAAKTRATTGA AATATTTTTWTTTGCCCCCCCTTGGAGTCAAGAAGGGTTTTTAGTTTTATCTTCTTYTTCTATTGAAGTTAAAAAAA

Figure 1B

Analysis of 26583 (537 aa)

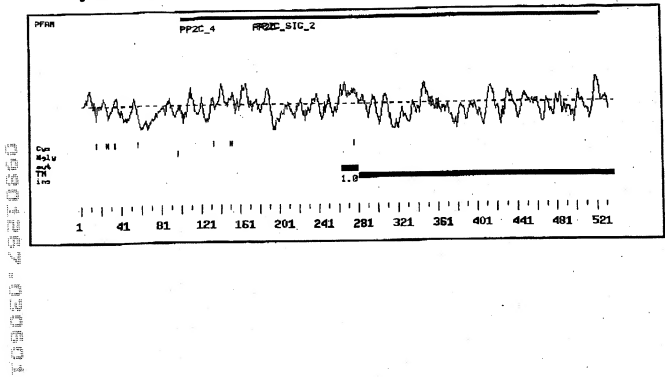


Figure 2

PP2C:	domain 1	o£	1, from 173 to 461: score 261.3, E = 1.3e-74 >ldvgvszmqgwrksmepahialkninssssgkdswsffavEDGhgGs 1 ++++ + F +++ ++++++ ++ g++ g+ +f ++++		SEQ. ID	NO:4
	26583	173		218	SEQ ID	NO:2
			qaakyagkhihk.tilaerksfpegdpwEmklsdledalkesfleadtde + + + + + + + + + + + + + + + + + + +			
	26583	219	Lidlntgestdidvkealinapkrldndisleaqvgdphsplny	262		
			elrsaeasaankvltkedlssGsTAvvalirgnkLyVANvGDSRavLcrn +++			
	26583	263	LVLRVAFSGATACVAHVDGVDLHVANTGDSRAMLGVQ	299		
	26583	300	gnaikw.avtltedhkpsnedererieaaggfvarvsngrvngvlav +++W+avtL++DH+++ne+E+eR++ ++++ + + + ++R++G L++ EEDGSWsAVTLSNDHNAQNERELERLKLEHPKSEAKSvvkQDRLLGLLMP		•	
	26583	350	SRAfGDielkpgsklgpees.l.e.a.ny.eyiks.peqlvtaer RAfGD+++K+ +1+++ +++++++ n++ey+k+ p++ +++++ taer FRAFGDVKFKWSIDLQKRVIeSgPdQlNDnexTKFiPPnyhtpPYLTAER	399		· .
	26583	400	dvtsstdltpdkDeflitAcDGLWDvvsdqevvdivfselsdgnksaedp +vt +++1+p+ D+F1+LA+DGLW++++ q+vv iv + 1+ + + EVT-YHRLRPQ-DKFLVLATDGLWETHHRQDVVRIVGEYLTGMH			
	26583	442	meaaeklvdesiargseDni<-* ++ ++ +			

Figure 3A

PP2C_4:	domain	1 0	f 1, from 99 to 523: score 338.5, E = 7.64-98					
			*->es.sgknlglryglgessmqgwrkpmEDahvirpffgvED p gkh +++ g+ +s++ +++ p+ED+ ++ + ++++ + gvfb		SEQ	ΙD	NO:5	
	26583	99		144	SEO	ID	NO: 2	
*	4000		2 W B De contract Shelled DE LEGIST & With contract and an American					
			GHGGseackflskniheilaeelsfdkdeslkens.e.lk.d.ep					
	•		GH+G ++++ +#++1+ ++a +1 +++ ++ an+ ar++ + ++ ++					
· · ·	26583	145	GHAGCACSQAVSERLFYYIAVSLLPHETLLEIENAVE:GRaL1PI1qwhk	194				
			gdksledvecalrkaflrtd					
			++++ +++ ++ ++ ++++++ +ln+g++++ dv+eal++aP+F+d					
3	26583	195	hpndyfskeasklyfnsl;TYWqElidLNtGESTDIDVKEALINAPKRLD	244				
			eei					
			++i+ + + +++++ + + +++TA+VA+++g +1+VAN+GDSR4					
2	6583	245	NDIsleadygdpnsflnylvlrvafagATACVAHVDGVDLHVANTGDSRA	294				
			vLcrngkd.swegvrtysavqLteDHkpanedEreRIeaaGCevepidre					
			+L+ + +d+9W sav L++DH++ ne+E+eR++ ++++ & +++					
•	26583	295	mlgvqeedgswsavtlsndhnaqnerelerlklehpkseaks	336				
			fvsngggvvvRVnGvvisLavsRalGDfelKk.ked.e.lieen.					
			+V ++ R++G L++ Ra+GD++-K+++++++++++++++++++++++++++++++++					
	26583	337	vvkqDRliglimpfrafgdvkfkmsidlqkfviEsgpdqlnd	378				
			rlyekfdprlpgkepyvsaePevtvvelsqtlvptedddfliLASDCLMD					
			++y+kf p+ ++++py++aePeut+++1 +++d+fl+LA+DGLM+					
7	6583	379	NEYTKPIPPNYHTPPYLTAEPEVTYHRLRPQDKFLVLATDGLWE	422				
			vlsnqeavdivrkhlrkgddk.evksaaqela.r.a.ds					
			+ q++V iV + 1++++++++++ +++++ +++++ ++					
7	26583	423	THUR ODVVRIVGEYLTCHOHIQ QPIAVGCYKVTlGqMGC11teRithmsEV	472				
			l.rskkhndpkesaklLvdlalkD#iTvvv					
			++++ ++1 X+ +++++ ++++ ++++ +++++++++++					
:	26583	473	fedgmaathLiRhaVGNNZFGTVDHERLSRMLSlpeelarmyRDDITIIV	522				
	٠.		V<-*					
	•		v					

Figure 3B

26583 Expression in Clinical Tissue Samples

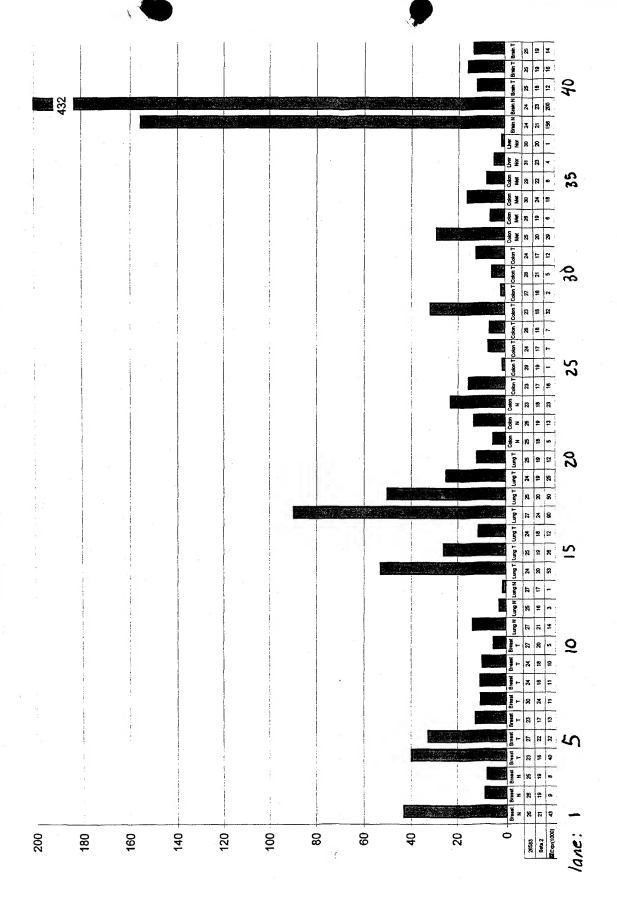


Figure 4